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## Cutis laxa-Marfanoid syndrome

**INSERM** 

## Source

INSERM. (1999). Orphanet: an online rare disease and orphan drug data base. <u>Cutis laxa-Marfanoid syndrome</u>. ORPHA:171719

A rare, genetic, developmental defect with connective tissue involvement syndrome characterized by neonatal cutis laxa, marfanoid habitus with arachnodactyly, pulmonary emphysema, cardiac anomalies, and diaphragmatic hernia. Mild contractures of the elbows, hips, and knees, with bilateral hip dislocation may also be associated. There have been no further descriptions in the literature since 1991.

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